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# Whole Genome Sequencing Services User Guide



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## Revision History

Part #	Revision	Date	Description of Change
15040892	D	June 2015	<ul> <li>Revised documentation to reflect changes in version 4 of the Illumina FastTrack WGS pipeline.</li> <li>Renamed Manta and Canvas to Isaac Structural Variant Caller and Isaac Copy Number Variant Caller, respectively.</li> </ul>
15040892	С	July 2014	Revised documentation to reflect changes in version 3 of the Illumina FastTrack WGS pipeline.
15040892	В	July 2013	Added Circos plot legend plus minor modifications.
15040892	A	April 2013	Initial release.



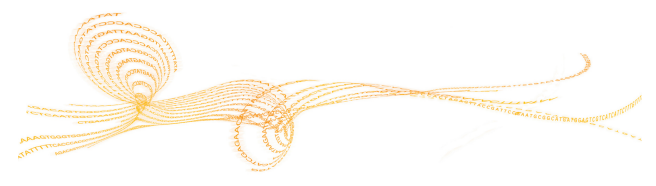
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# Getting Started

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## Whole Genome Sequencing Service

The Whole Human Genome Sequencing Service Informatics Pipeline leverages a suite of proven algorithms to detect genomic variants comprehensively and accurately. High-quality sequence reads are aligned using the Isaac Aligner (see *Isaac Aligner* on page 26 for details). Variant calling is performed using the Isaac Variant Caller (see *Isaac Variant Caller* on page 28 for details). Two complementary approaches enable detection of large structural variations:

- Read depth analysis by Isaac Copy Number Variant Caller. See *Isaac Copy Number Variant Caller* on page 35.
- Discordant paired-end analysis by Isaac Structural Variant Caller (Manta). See *Isaac Structural Variant Caller* on page 40.

Identified variants are then annotated and compiled into a summary PDF.

This document provides an overview of the source and contents of the main files that Illumina creates using this informatics pipeline. This document also provides information about key algorithms, such as the Isaac Variant Caller, Isaac SV Caller, and Isaac CNV Caller. The aim of this document is to help you understand the Whole Genome Sequencing data package that you receive from Illumina.

The following versions of software packages are utilized in the Control Software (CS) v4.0.2 pipeline.

Software	Version	Purpose
Isaac Aligner	6.15.01	Align reads to the human hg19 reference.
Isaac Variant Caller	2.1.4	Germline SNV and indel caller.
Isaac Copy Number Variant Caller	1.1.0	Germline copy number variant caller.
Isaac Structural Variant Caller	0.23.1	Germline and somatic structural variant caller.

## **Data Delivery**

Illumina FTS currently provides data delivery through the following choices.

## Illumina Hard Drive Data Delivery

Illumina FastTrack Services ships data on 1 or more hard drives. The hard drives are formatted with the NTFS file system and can optionally be encrypted.

The data on the hard drive are organized in a folder structure with 1 top-level folder per sample or analysis.

## Illumina Cloud Data Delivery

Illumina FastTrackServices uploads data to a cloud container. Illumina currently supports uploads to the Amazon S3 service. Upload data are organized per upload batch by date under an Illumina\_FTS prefix. For example, a sample in a batch uploaded on February 1, 2014 would be found with the prefix Illumina\_FTS/20140201/SAMPLE\_BARCODE in the container. Contact your FastTrack Services project manager to enable cloud delivery.

# Analysis Deliverables

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## Analysis Folder Structure Overview

This section details the files and folder structure for the single whole genome deliverable. Several folders are batched together at delivery but each sample follows the same underlying format.

The files and folders generated for the whole genome deliverable are all keyed off a unique sample identifier [Sample\_Barcode]. Usually, this unique identifier is the barcode associated with a sample in the lab (eg, LP6000001-DNA\_A01) but can be a common sample identifier for reference samples (eg, NA12878).

## Result Folder Structure

Under each Sample folder, you can find the following file structure that contains analysis results.

## [Sample\_Barcode] Assembly ■ [Sample Barcode].bam — Archival \*.bam file for sample. ■ [Sample Barcode].bam.bai – Index for \*.bam file ☐ [Sample\_Barcode].SummaryReport.csv—Summary report in \*.csv format [Sample\_Barcode].SummaryReport.pdf—Summary report in \*.pdf format Genotyping [Sample\_Barcode] idats—Folder containing genotyping intensity data files for the sample (\*.idat files) and genotyping sample sheet. [Sample\_Barcode].Genotyping.vcf.gz—Genotyping SNPs mapped to reference in \*.vcf format. □ [Sample\_Barcode].GenotypingReport.txt—Genotyping SNPs tab delimited report. **T**Variations [Sample\_Barcode].CNV.vcf.gz—Copy number calls (10 kb +) in \*.vcf format. ☐ [Sample Barcode].Indels.vcf.gz—Small Insertion/Deletion calls (1bp-50 bp) in \*.vcf format. □ [Sample\_Barcode].SNPs.vcf.gz—Single nucleotide polymorphism (SNVs) calls in \*.vcf format. [Sample\_Barcode].SV.vcf.gz—Large Structural Variation calls (51 bp-10 kb) in \*.vcf format. [Sample\_Barcode].genome.vcf.gz—Genome \*.vcf file containing SNVs, indels, and reference covered regions [Sample\_Barcode].vcf.gz—\*.vcf file containing basic annotations and SNV

md5sum.txt—checksum file for confirming file consistency.



#### NOTE

and indel calls.

All the \*.vcf files that Illumina provides are compressed and indexed using tabix. For details about tabix, see the tabix manual in SAMtools (at samtools.sourceforge.net/tabix.shtml). The tabix index shows up as an additional [Sample\_Barcode].TYPE.vcf.gz.tbi file. It can be used for fast retrieval of targeted regions in the associated vcf.gz file



#### NOTE

For some VCF files, a binary format of the annotations and their indexes are contained in corresponding .vcf.ant and .vcf.ant.idx files respectively. If the .vcf.ant file is maintained in the same directory as its VCF file, the annotation information can be visualized alongside the variant call information when imported to VariantStudio.

## Assembly

The assembly folder contains the sequence data used to assemble the sample genome.

## [Sample\_Barcode].bam

The included archival BAM file contains all pass filter reads input into the analysis pipeline for a sample and includes aligned, duplicate, and unaligned reads. To reduce the data storage footprint while not compromising accuracy, Illumina has reduced quality score resolution in BAM files. In practice, this means that the more commonly used 40+ possible Q-scores have been reduced to 8 bins. This transformation is performed on instrument and is calibrated for each individual quality table.

For details about how Illumina has reduced storage requirements while maintaining compatibility and accuracy, see the Reducing Whole-Genome Data Storage Footprint white paper on the Illumina product literature page.

## [Sample\_Barcode].bam.bai

This is the SAMtools BAM index for the BAM file. This file can be used with SAMtools and other tools utilizing the SAMtools specification for fast retrieval of targeted regions in the associated BAM file.

#### **BAM File Details**

The included BAM file adheres to the SAM format specification wherever possible. The following sections cover BAM file details that are not evident in the specification:

- ▶ Singleton / Shadow Pairs
- Read Groups: RG
- ▶ Read name: RNAME
- ▶ Bitwise Flag Notes: FLAG
- Extended Tags / Optional Fields
- MAPO

## Singleton / Shadow Pairs

Singleton/shadow pairs refer to pairs for which the aligner was unable to determine the alignment of 1 of the ends. The determined end is the singleton and the undetermined end is the shadow. Shadows are assigned the position of the end that does align. To maintain SAMtools format compatibility, the shadows are stored in the BAM file immediately after their respective singletons, with CIGAR empty and corresponding flag (4) set. Shadows can be retrieved using the following SAMtools command:

```
samtools view -f 4 input.bam > output.sam
```

## Read Groups: RG

Where possible, unique flow cell-lane-index mappings split up the read groups in the BAM. The following is an example from a BAM header:

```
@RG ID:0 PL:ILLUMINA SM:NA12878 PU:C0L0AACXX:1:none
@RG ID:1 PL:ILLUMINA SM:NA12878 PU:C0L54ACXX:7:none
@RG ID:2 PL:ILLUMINA SM:NA12878 PU:C0L54ACXX:8:none
```

In the example, the read group 0 is derived from the flow cell barcode ID C0L0AACXX, lane 1, without a specified index for sample NA12891. In this example, read groups 1 and 2 are from a different flow cell C0L54ACXX, lanes 7 and 8.

#### Read name: RNAME

The read name consists of the following pattern detailing the flow cell, lane, and tile on which the sample was run:

```
flowcell-id ":" lane-number ":" tile-number ":"cluster-id
    ":"cluster-id-alt"
```

Table 1 RNAME Variables

ID	Description
cluster-id	Unpadded 0-based cluster id in the order in which the clusters appear within the tile.
flowcell-id	Flow cell barcode.
cluster- id-alt	In cases where the x:y coordinates from the flow cell were preserved, this column will contain the y-coordinate, while the cluster-id will contain the x-coordinate. Otherwise this will always contain "0".
lane-number	Lane number 1–8.
tile-number	Unpadded tile number.

## Bitwise Flag Notes: FLAG

The bitwise flags used are as follows.

Bit	Description	Note
0x1	Template having multiple segments in	Always set. (on for paired reads)
OXI	sequencing.	Tiways set. (on for panea reads)
0x2	Each segment properly aligned according to	Pair matches dominant template
	the aligner.	orientation.
0x4	Segment unmapped. Set for unmapped reads.	
0x8	Next segment in the template unmapped.	Paired read is unmapped.
0x10	SEQ being reverse complemented. Read mapped to—strand of refere	
0x20	SEQ of the next segment in the template being	Paired read mapped to -strand of
	reversed.	reference.
0x40	The first segment in the template.	Read 1 sequence.
0x80	The last segment in the template.	Read 2 sequence.
0x100	Secondary alignment.	Isaac does not produce secondary
		alignments.
0x200	Not passing quality controls.	Nonpass filter reads are not included
		(always off).
0x400	PCR or optical duplicate.	Read 1 and Read 2 were marked as
		duplicate reads.

## **Extended Tags and Optional Fields**

The aligner produces the following fields in the BAM file.

#### Table 3 BAM File Fields

Field	Description
AS	Pair alignment score.
ВС	Barcode string.
NM	Edit distance (mismatches and gaps) including the soft-clipped parts of the read.
OC	Original CIGAR for the realigned reads.
RG	Isaac read groups correspond to unique flow cell-lane-barcodes.
SM	Single read alignment score.

## Mapping Quality (MAPQ)

For pairs that match the dominant template orientation, the MAPQ value in the AS field is capped. For reads that are not members of a pair matching the dominant template orientation, the MAPQ value in the SM field is capped at 60. The MAPQ could be downgraded to 0 or set to be unknown (255) for alignments that do not have enough evidence to be correctly scored.

## Genotyping

If available, variants called using the Infinium platform are compared to sequencing calls to confirm identity and make sure that data are of high quality. This folder contains the results of the genotyping SNP calls and the necessary files needed to regenerate them.

To download the end-user documentation for the GenomeStudio Genotyping Module, go to support.illumina.com/documents/MyIllumina/d2c2c169-36c7-4613-89d6-bf34588a7624/GenomeStudio GT Module v1.0 UG 11319113 RevA.pdf.

## [Sample\_Barcode]\_idats

This folder contains the GRN.idat and RED.idat intensity files and the sample sheet for a genotyping sample. These files along with the manifest, cluster, and genotyping product files can be imported into the Illumina GenomeStudio software genotyping module (www.illumina.com/software/genomestudio\_software.ilmn) to reproduce the genotyping calls. The genotyping product files can be found on the Array support page in the Downloads tab. To find the version of array chip used for your project, refer to the sample sheet in each sample folder. The sample sheet can be found in the following directory of each sample folder: Sample\_Barcode\Genotyping\Sample\_Barcode\_idats If available, \*.gtc files with genotype call files for use as input into genotyping software for reanalysis are also included.

## [Sample\_Barcode].Genotyping.vcf.gz

This file contains the genotyping SNPs in VCF format. The genotyping SNPs were mapped to the reference using megaBLAST and filtered in the following manner.

#### **Exclusions:**

- intensity only SNPs
- any match not aligning to the SNP
- any probe with a hamming distance greater than or equal to 5
- any probe where the highest scoring mapping site is not the best matching site (ie, there is another site or sites within an identical hamming distance)

Any genotyping probe not matching the reference or excluded from the mapping will be mapped to chromosome "NA" in the \*.vcf file.

The following fields are utilized in the \*.vcf file.

Table 4 INFO Fields

ID	Description
AL	Array alleles relative to the design strand of the array probe.
ST	The strand for the array alleles relative to the reference. A dash ( - ) denotes a reverse compliment.
GC	The GenCall score from the genotyping SNP call. (0.15 cut off applied by default).
GT	Genotype per VCF specification.

#### Table 5 FORMAT Fields

ID	Description
GC	The GenCall score from the genotyping SNP call. (0.15 cut off applied by default).
GT	Genotype per VCF specification.

#### Table 6 FILTER Fields

ID	Description
GTEX	The exclude genotype filter. The genotype was excluded in the mapping, possibly because the probe failed to find a reference map, failed to map uniquely, or was an intensity-only based probe.
NOCALL	Genotype value was not called on array.

## $[Sample\_Barcode]. Genotyping Report.txt$

This file contains the genotyping report that is output from the GenomeStudio Genotyping Module. Illumina provides the genotyping report as a tab-delimited text file and includes a header followed by at least the following columns.

Table 7 Genotyping Report Columns

Column	Description
Allele1 – Design	The A allele call that is relative to the probe.
Allele1 - Forward	The A allele call that is relative to the submitted sequence.
Allele2 - Design	The B allele call that is relative to the probe.
Allele2 – Forward	The B allele call that is relative to the submitted sequence.
GC Score	The GenCall score. This score is a quality metric assigned to every genotype called, and generally indicates their reliability. GC scores have a maximum of 1, and are calculated using information from the clustering of the samples. Each SNP is evaluated based on the angle of the clusters, dispersion of the clusters, overlap between clusters, and intensity. Genotypes with lower GC scores are located furthest from the center of a cluster and have a lower reliability.
Sample Barcode	The internal process identifier.
SNP Name	The SNP identifier. An rsID for dbSNP content.

### **Variations**

The variations folder contains the variant call output in VCF 4.1 format for the sample. Each variant file that Illumina provides is compressed and includes an index that was created using tabix, for fast range-based access. This is a summary of the outputs for each sample. See *Introduction* on page 24 for details. The VCF files are annotated with the FastTrack Services Annotation Pipeline. See *Illumina FastTrack Services Annotation Pipeline* on page 46 for details.

## [Sample\_Barcode].CNV.vcf.gz

The CNV file contains large copy number variants from 10 kb+ output from the Isaac CNV Caller. The following fields are utilized in the VCF file.

Table 8 INFO Fields

ID	Description
END	End position of the variant described in this record.
SVTYPE	Type of structural variant.

#### Table 9 ALT Fields

ID	Description
CNV	Copy number variable region.

#### Table 10 FORMAT Field

ID	Description
ВС	Number of bins in the region.
CN	Copy number genotype for imprecise events.
GT	Genotype.
RC	Mean counts per bin in the region.

#### Table 11 FILTER Fields

ID	Description
q10	Quality below 10.
L10kb	For a small variant (< 1000 base), the fraction of reads with MAPQ=0 around either break-end that exceeds 0.4.

## [Sample\_Barcode].SNPs.vcf.gz and [Sample\_Barcode].Indels.vcf.gz

### [Sample\_Barcode].SNPs.vcf.gz and [Sample\_Barcode].Indels.vcf.gz

The SNV and indel files list the single nucleotide polymorphisms and indels (respectively) that were called by the Isaac Variant Caller. Small indels are limited to 50 bp. The VCF file contains the following fields.

Table 12 INFO Fields

ID	Description
CIGAR	The CIGAR alignment for each alternate indel allele.
END	The end position of the region described in this record.
RU	The smallest repeating sequence unit extended or contracted in the indel allele relative to the reference. RUs are not reported if longer than 20 bases.
IDREP	Number of times RU is repeated in an indel allele.
REFREP	Number of times RU is repeated in the reference.
SNVHPOL	SNV contextual homopolymer length.
SNVSB	SNV site strand bias.

Table 13 FORMAT Fields

Table 15 FORWAT FIELDS		
ID	Description	
AD	Allelic depths for the ref and alt alleles in the order listed. For indels, this value includes only reads that confidently support each allele. Specifically, includes reads for which the posterior probability is 0.999 or higher that the read contains an indicated allele versus all other intersecting indel alleles.	
DP	Filtered base call depth used for site genotyping.	
DPF	Base calls filtered from input before site genotyping.	
DPI	Read depth associated with indel, taken from the site preceding the indel.	
GQ	Genotype quality.	
GQX	Minimum Phred genotype quality. Annotated as {Genotype quality assuming variant position, Genotype quality assuming nonvariant position}.	
GT	Genotype.	
OPL	Original Phred-scaled genotype likelihood (PL) value before ploidy correction. Only applies to sites with "HAPLOID_CONFLICT" FILTER applied.	

Table 14 FILTER Fields

ID	Description
GENDER_ CONFLICT	Genotype is inconsistent with sample gender.
HAPLOID_ CONFLICT	Locus has heterozygous genotype in a haploid region.
HighDepth	The locus depth is greater than 3x the mean chromosome depth.

ID	Description
HighDPFRatio	The fraction of base calls filtered out at a site is greater than 0.3.
HighSNVSB	SNV strand bias value (SNVSB) exceeds 10.
IndelConflict	The locus is in a region with conflicting indel calls.
LowGQX	Locus GQX is less than 30 or not present.
SiteConflict	The site genotype conflicts with the proximal indel call. This is typically a heterozygous SNV call made inside a heterozygous deletion.

## [Sample\_Barcode].SV.vcf.gz

The SV file contains structural variants from  $50\ bp-10\ kb$  called within the sample by the Isaac SV Caller. The VCF file contains the following fields.

Table 15 ALT Fields

ID	Description
BND	Translocation break-end.
COMPLEX	Unknown Candidate Type.
DEL	Deletion.
DUP:TANDEM	Tandem Duplication.
INS	Insertion.
INV	Inversion.

Table 16 INFO Fields

ID	Description
BND_DEPTH	Read depth at local translocation break-end.
BND_PAIR_COUNT	Confidently mapped reads supporting this variant at this break-end (it is possible that mapping is not confident at remote break-end).
CIEND	Confidence interval around END.
CIGAR	CIGAR alignment for each alternate indel allele.
CIPOS	Confidence interval around POS.
DOWNSTREAM_ PAIR_COUNT	Confidently mapped reads supporting this variant at this downstream break-end (it is possible that mapping is not confident at upstream break-end).
END	End position of the variant described in this record.
HOMLEN	Length of base pair identical micro-homology at event breakpoints.

ID	Description
HOMSEQ	Sequence of base pair identical micro-homology at event breakpoints.
IMPRECISE	Imprecise structural variation.
MATE_BND_DEPTH	Read depth at remote translocation mate break-end.
MATEID	ID of mate break-end .
PAIR_COUNT	Read pairs supporting this variant where both reads are confidently mapped.
SVINSLEN	Length of microinsertion at event breakpoints.
SVINSSEQ	Sequence of microinsertion at event breakpoints.
SVLEN	Difference in length between REF and ALT alleles.
SVTYPE	Type of structural variant described in the ALT field.
UPSTREAM	Reference sequence upstream of the variant.
UPSTREAM_PAIR_ COUNT	Confidently mapped reads supporting this variant at the upstream break-end (it is possible that mapping is not confident at downstream break-end).

#### Table 17 FORMAT Fields

ID	Description
GQ	Genotype Quality.
GT	Genotype.
PR	Spanning paired read support for the REF and ALT alleles in the order listed.
SR	Split reads for the REF and ALT alleles in the order listed, for reads where P (allele $\mid$ read) > 0.999.

## [Sample\_Barcode].genome.vcf.gz

The genome \*.vcf file contains \*.vcf formatted output for the SNVs, indels and block compressed nonvariant position output. You can use this file to compare variants and covered regions between samples quickly. The filters and INFO fields are a combination of both the SNV and indel \*.vcf files listed below along with the block compressed specific flags. See *Genome VCF* (*gVCF*) on page 30 for details. For additional INFO fields pertaining to annotation information, see *Introduction* on page 24.

## [Sample\_Barcode].vcf.gz

This VCF file contains SNV and indel calls, along with basic annotations. Nonvariant positions are not included.

Table 18 INFO Fields

ID	Description		
AA	The inferred allele ancestral to the chimpanzee/human lineage.		
CF1000G	The allele frequency from all populations of 1000 genomes data.		
BLOCKAVG_min30p3a	Nonvariant site block. All sites in a block are constrained to be nonvariant, have the same filter value, and have all sample values in range [x,y], $y \le \max(x+3,(x*1.3))$ . All printed site block sample values are the minimum observed in the region spanned by the block		
CIGAR	The CIGAR alignment for each alternate indel allele.		
CLINVAR	Clinical significance.		
CSQT	Transcript consequence as predicted by VEP version 72 using transcripts from Ensmbl.		
	Annotated as: HGNC   TranscriptID   Consequence		
CSQR	Regulatory consequence type as predicted by VEP version 72 using features from Ensmbl.		
	Annotated as: RegulatoryID Consequence		
COSMIC	The numeric identifier for the variant in the Catalogue of Somatic Mutations in Cancer (COSMIC) database.		
END	The end position of the region described in this record		
EVS	Allele frequency, sample count, and coverage taken from the Exome Variant Server (EVS).		
	Annotated as: AlleleFreqEVS EVSCoverage EVSSamples		
GMAF	Global minor allele frequency (GMAF); technically, the frequency of the second most frequent allele.		
	Annotated as: GlobalMinorAllele   AlleleFreqGlobalMinor		
IDREP	Number of times RU is repeated in an indel allele.		
phastCons	Denotes if the variant is an identical or similar sequence that occurs between species and maintained between species throughout evolution		
REFREP	Number of times RU is repeated in the reference.		
RU	The smallest repeating sequence unit extended or contracted in the indel allele relative to the reference. RUs are not reported if longer than 20 bases.		
SNVHPOL	SNV contextual homopolymer length.		
SNVSB	SNV site strand bias.		

Table 19 FORMAT Fields

ID	Description			
AD	Allelic depths for the ref and alt alleles in the order listed. For indels, this value includes only reads that confidently support each allele. Specifically, includes reads for which the posterior probability is 0.999 or higher that the read contains an indicated allele versus all other intersecting indel alleles.			
DP	Filtered base call depth used for site genotyping.			
DPF	Base calls filtered from input before site genotyping.			
DPI	Read depth associated with indel, taken from the site preceding the indel.			
GQ	Genotype quality.			
GQX	Minimum Phred genotype quality. Annotated as {Genotype quality assuming variant position, Genotype quality assuming nonvariant position}.			
GT	Genotype.			

Table 20 FILTER Fields

ID	Description	
HighDepth	The locus depth is greater than 3x the mean chromosome depth.	
HighDPFRatio	The fraction of base calls filtered out at a site is greater than 0.3.	
HighSNVSB	SNV strand bias value (SNVSB) exceeds 10.	
IndelConflict	The locus is in a region with conflicting indel calls.	
LowGQX	Locus GQX is less than 30 or not present.	
SiteConflict	The site genotype conflicts with the proximal indel call. This is typically a heterozygous SNV call made inside a heterozygous deletion.	

## Summary Report

The [Sample\_Barcode].SummaryReport.pdf report contains an overview of the results for the sample. In the report you will find the following:

- ▶ Sample Information
- Library Specifications
- Data Volume
- ▶ Passing Filter and Aligned Base call Quality Score Distribution
- Coverage Summary
- Non-N Reference Coverage Distribution
- ▶ SNV / Indel Assessment
- Variant Statistics
- Structural Variants Summary

## Sample Information

This section contains information associated with the sample from the included sample manifest.

## Library Specifications

This section describes details related to the library prep used in the sample.

Table 21 Library Specification Values

Value	Description	
Fragment Length Median	Median fragment length of library sequence fragments calculated as for each pair of mapped reads. For normal reads, this value includes both reads, along with the unsequenced insert between the reads.	
Fragment Length SD	The standard deviation of fragment lengths around the median.	
Read Length	Read lengths used in the build.	
Read Type	Will be pared end for the standard Whole Genome Sequencing workflow.	

#### **Data Volume**

This table in the reports the volume of data input into the assembly process (and in the associated BAM file).

Table 22 Data Volume Table Values

Value	Description	
Passing Filter	Yield is the number of gigabases of data (PASS filter data only) input into the build.  % Bases ≥ Q30 — where Q30 is the percent of the sequence data that has a Q-score of Q30 or greater.  Note: Q-score binning does not affect this measure.	
Passing Filter and Aligned	Same as passing filter but reported only for the subset of data that aligns to the genome.	

## Passing Filter and Aligned Base call Quality Score Distribution

This table details the quality score distribution for the aligned reads for a sample. The spiky appearance of the graph is due to the effect of quality score binning.

## Coverage Summary

The coverage summary reports the distribution of depth of coverage across the genome. Coverage is calculated from bases not flagged as duplicates and for which both read pairs map unambiguously.

Table 23 Coverage Summary Values

Value	Description	
%≥ 5/10/20x coverage	Number of non-N reference autosomal positions that have greater or equal to 5/10/20 fold coverage.	
% The percent of autosomal non-N reference genome in gVCF file with a PAS status.		
Average Coverage	Mean coverage across the genome defined as "bases used in variant calling over autosomal regions" / "total non-N reference length of autosomal regions".	

## Non-N Reference Coverage Distribution

This histogram of coverage depth uses the same definition of coverage as the Coverage Summary.

### SNV and Indel Assessment

These tables provide the total number of SNVs and Indels overlapping known variants and genes, exons and coding regions. All counts only use PASS filter variants where applicable.

Table 24 SNV and Indel Assessment Table Values

Value	Description	
% Array Agreement	The percentage of concordant SNVs between the genotyping and sequencing SNVs.  Note: Only PASS filter SNVs are compared.	
% in Coding	Percent of PASS filter variants overlapping a coding position for any annotated transcript.	
% in dbSNP	Percent of PASS filter variants that overlap a dbSNP identifier in annotation.	
% in Exons	Percent of PASS filter variants overlapping a coding, 5' UTR or 3' UTR position for any annotated transcript.	
% in Genes	Percent of PASS filter variants overlapping a coding, 5' UTR, 3' UTR or intron position for any annotated transcript.	
Het/Hom	The ratio of heterozygous to homozygous PASS filter variants reported.	

Value	Description
Ti/Tv	The transition/tranversion ratio for reported variants relative to the reference base or bases.
Total	Total number of PASS filter SNVs reported.

#### Variant Statistics

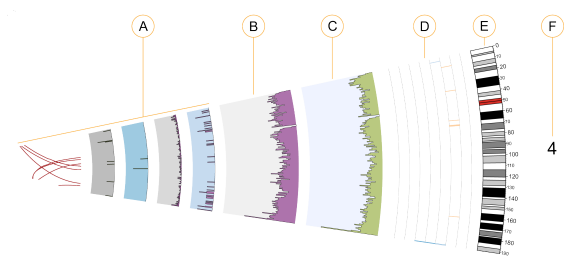
This table breaks down SNVs and indels into total counts in overlapping regions and annotated consequences. Complex indels are split into deletions and insertions where appropriate. Consequence types for overlapping transcripts are counted under the most severe transcript consequence according to the annotation.

## Structural Variants Summary

This table breaks CNV and SV output into the classes of variants called. Their total PASS count and the number of overlapping genes are based on the annotation pipeline (see *Illumina FastTrack Services Annotation Pipeline* on page 46).

#### Circos Plot of Genome Variations

The Circos plot provides visualization of structural variation, ploidy, and structural variations reported in the genome variation files (VCF). The Circos plot displays genome variation data in tracks with chromosomes circularly arranged. Following is an example legend. Labels are described from inside the circle to the outside.



	Circos Plot Legend		
Legend	Label (From Inner Circle to Outer Circle)	Description	
A	Structural variants	The structural variants described in [Sample_ Barcode].SV.vcf.gz are plotted in the central portions of the Circos plot. From inner to outer (left to right in the legend):	
		Red Links—Translocation break-ends	
		Dark Gray — Tandem Duplications per Mb	
		Dark Blue – Inversions per Mb	
		Grey — Deletions per Mb	
		Blue — Insertions per Mb	
B Number indels per Mb		The density of PASS indels reported in [Sample_Barcode].SV.vcf.gz in 1 Mb windows.	
		The scale of Y-axis in the histogram indicates the counts.	
С	Number of SNVs per Mb	The density of PASS SNVs reported in [Sample_Barcode].SV.vcf.gz 1 Mb windows, arbitrarily scaled in a histogram with Y-axis pointing inward.	
D	Copy number variation	The copy number variations from [Sample_Barcode].CNA.vcf.gz file. The scale of Y-axis in the histogram indicates the called level.	
		• Orange bar—loss of copy (fewer than 2 copies)	
		• Blue bar – gain of copy (greater than 2 copies, max of 5)	
Е	Karyotype/Chromosome position	The standard Circos ideogram defining the chromosome position, identity, and color of cytogenetic bands and the	
	position	reference coordinates along the chromosome.	

## Data Integrity

The md5sum.txt file is provided as a means of checking the integrity of the sample files and folders. Immediately after sample quality check, the md5sums, or compact digital fingerprint, for every file in the directory tree are generated. If media failures compromise data integrity, you can use the md5sum tool to find the inconsistencies. Use the tool to compare the hash from the provided md5sum file to one generated from the downloaded file.

On a Unix system, you can use the following commands to perform an md5sum check (assuming the utility is installed):

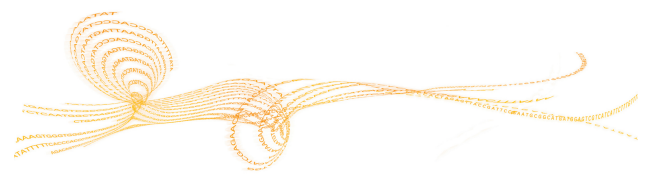
- % cd [Sample\_Barcode]
- ▶ % md5sum –c md5sum.txt

The check verifies every file and require approximately 30–45 minutes to complete. Any errors are listed in the output.

In Windows, there are various command line and GUI tools available to perform an md5sum check. The Cygwin tools provide a utility identical to Linux.

# Analysis Overview

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Isaac Structural Variant Caller	



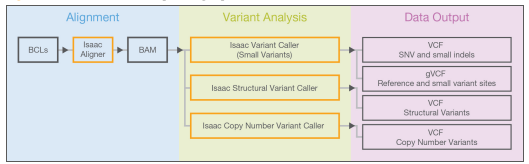
## Introduction

After the sequencer generates base calls and quality scores, the resulting data are analyzed in 2 steps—alignment to the reference genome followed by assembly and variant calling.

Alignment and variant calling are performed with the Isaac Alignment Software, Isaac Variant Caller, Isaac CNV Caller, and Isaac SV Caller. The following output is produced:

- ▶ Realigned and duplicate marked reads in a \*.bam file format.
- Variants in a VCF file format.
- ▶ An additional Genome VCF (gVCF) file. This file features an entry for every base in the reference, which differentiates reference calls and no calls, and a summary of quality. The reference calls are block compressed and all single nucleotide polymorphisms and indels are included. Currently Structural Variants and CNVs are kept in separate files.

Figure 1 Whole-Genome Sequencing Pipeline



## Genome Specific Details

Illumina currently uses hg19 from UCSC as a reference genome. The chromosome naming scheme follows the UCSC conventions of chr1-22, chrX, chrY, chrM. The pseudoautosomal region (PAR) of the Y chromosome is masked out with N's. The result of this is that any mappings occurring in the PAR region map to the X chromosome. Currently, only the main chromosomes and mitochondria are used in the reference; none of the nonmapped contigs are included. As per GATK specification for UCSC, chrM is the first chromosome followed by the rest in karyotypic order.

The hg19 PAR regions are defined as follows.

Table 26 hg19 PAR regions

Name	Chr	Start	Stop
PAR#1	X	60,001	2,699,520
PAR#2	X	154,931,044	155,260,560
PAR#1	Y	10,001	2,649,520
PAR#2	Y	59,034,050	59,363,566

You can find links to Illumina iGenomes references here: support.illumina.com/sequencing/sequencing\_software/igenome.ilmn.



NOTE

The version of hg19 provided in iGenomes is not PAR masked.

## Isaac Aligner

The Isaac Aligner<sup>1</sup> aligns DNA sequencing data, single or paired-end, with read lengths of 32–150 bp and low error rates using the following steps:

- ▶ **Candidate mapping positions**—Identifies the complete set of relevant candidate mapping positions using a 32-mer seed-based search.
- ▶ **Mapping selection**—Selects the best mapping among all candidates.
- ▶ **Alignment score** Determines alignment scores for the selected candidates based on a Bayesian model.
- ▶ **Alignment output**—Generates final output in a sorted duplicate-marked BAM file, and summary file.
- Come Raczy, Roman Petrovski, Christopher T. Saunders, Ilya Chorny, Semyon Kruglyak, Elliott H. Margulies, Han-Yu Chuang, Morten Källberg, Swathi A. Kumar, Arnold Liao, Kristina M. Little, Michael P. Strömberg and Stephen W. Tanner (2013) Isaac: Ultra-fast whole genome secondary analysis on Illumina sequencing platforms. Bioinformatics 29(16):2041-3 bioinformatics.oxfordjournals.org/content/29/16/2041

## Candidate Mapping

To align reads, the Isaac Aligner first identifies a small but complete set of relevant candidate mapping positions. The Isaac Aligner begins with a seed-based search using 32-mers from the extremities of the read as seeds. Isaac Aligner performs another search using different seeds for only those reads that were not mapped unambiguously with the first pass seeds.

## Mapping Selection

Following a seed-based search, the Isaac Aligner selects the best mapping among all the candidates. For paired-end data sets, all mappings where only one end is aligned (called orphan mappings) trigger a local search to find additional mapping candidates. These candidates (called shadow mappings) are defined through the expected minimum and maximum insert size. After optional trimming of low quality 3' ends and adapter sequences, the possible mapping positions of each fragment are compared. This step takes into account pair-end information (when available), possible gaps using a banded Smith-Waterman gap aligner, and possible shadows. The selection is based on the Smith-Waterman score and on the log-probability of each mapping.

## Alignment Scores

The alignment scores of each read pair are based on a Bayesian model, where the probability of each mapping is inferred from the base qualities and the positions of the mismatches. The final mapping quality (MAPQ) is the alignment score, truncated to 60 for scores above 60, and corrected based on known ambiguities in the reference flagged during candidate mapping. Following alignment, reads are sorted. Further analysis is performed to identify duplicates and optionally to realign indels.

## **Alignment Output**

After sorting the reads, the Isaac Aligner generates compressed binary alignment output files, called BAM (\*.bam) files, using the following process:

- Marking duplicates—Detection of duplicates is based on the location and observed length of each fragment. The Isaac Aligner identifies and marks duplicates even when they appear on oversized fragments or chimeric fragments.
- ▶ **Realigning indels**—The Isaac Aligner tracks previously detected indels, over a window large enough for the current read length, and applies the known indels to all reads with mismatches.
- ▶ Generating BAM files—The first step in BAM file generation is creation of the BAM record, which contains all required information except the name of the read. The Isaac Aligner reads data from base call (BCL) files that were written during base calling on the sequencer to generate the read names. Data are then compressed into blocks of 64 kb or less to create the BAM file.

## Isaac Variant Caller

The Isaac Variant Caller identifies single nucleotide variants (SNVs) and small indels using the following steps:

- ▶ **Read filtering**—Filters out reads failing quality checks.
- ▶ **Indel calling**—Identifies a set of possible indel candidates and realigns all reads overlapping the candidates using a multiple sequence aligner.
- ▶ **SNV calling**—Computes the probability of each possible genotype given the aligned read data and a prior distribution of variation in the genome.
- ▶ Indel genotypes Calls indel genotypes and assigns probabilities.

#### Indel Candidates

Input reads are filtered by removing any of the following:

- ▶ Reads that failed base calling quality checks.
- Reads marked as PCR duplicates.
- Paired-end reads not marked as a proper pair.
- Reads with a mapping quality less than 20.

### **Indel Calling**

The variant caller proceeds with candidate indel discovery and generates alternate read alignments based on the candidate indels. As part of the realignment process, the variant caller selects a representative alignment to be used for site genotype calling and depth summarization by the SNV caller.

## **SNV Calling**

The variant caller runs a series of filters on the set of filtered and realigned reads for SNV calling without affecting indel calls. First, any contiguous trailing sequence of N base calls is trimmed from the ends of reads. Using a mismatch density filter, reads having an unexpectedly high number of disagreements with the reference are masked, as follows:

- ▶ The variant caller treats each insertion or deletion as a single mismatch.
- ▶ Base calls with more than 2 mismatches to the reference sequence within 20 bases of the call are ignored.
- ▶ If the call occurs within the first or last 20 bases of a read, the mismatch limit is applied to a 41-base window at the corresponding end of the read.
- ▶ The mismatch limit is applied to the entire read when the read length is 41 or shorter.

## **Indel Genotypes**

The variant caller filters out all bases marked by the mismatch density filter and any N base calls that remain after the end-trimming step. These filtered base calls are not used for site-genotyping but appear in the filtered base call counts in the variant caller output for each site.

All remaining base calls are used for site-genotyping. The genotyping method heuristically adjusts the joint error probability that is calculated from multiple observations of the same allele on each strand of the genome. This correction accounts for the possibility of error dependencies.

This method treats the highest-quality base call from each allele and strand as an independent observation and leaves the associated base call quality scores unmodified. Quality scores for subsequent base calls for each allele and strand are then adjusted. This adjustment is done to increase the joint error probability of the given allele above the error expected from independent base call observations.

## Variant Call Output

After the SNV and indel genotyping methods are complete, the variant caller applies a final set of heuristic filters to produce the final set of calls in the output.

The output in the genome variant call (gVCF) file captures the genotype at each position and the probability that the consensus call differs from reference. This score is expressed as a Phred-scaled quality score.

# Genome VCF (gVCF)

Human genome sequencing applications require sequencing information for both variant and nonvariant positions, yet there is no common exchange format for such data. gVCF addresses this issue.

gVCF is a set of conventions applied to the standard variant call format (VCF). These conventions allow representation of genotype, annotation, and additional information across all sites in the genome, in a reasonably compact format. Typical human wholegenome sequencing results expressed in gVCF with annotation are less than 1.7 GB, or about 1/50 the size of the BAM file used for variant calling.

gVCF is also equally appropriate for representing and compressing targeted sequencing results. Compression is achieved by joining contiguous nonvariant regions with similar properties into single 'block' VCF records. To maximize the utility of gVCF, especially for high stringency applications, the properties of the compressed blocks are conservative. Block properties such as depth and genotype quality reflect the minimum of any site in the block. The gVCF file is also a valid VCF v4.1 file, and can be indexed and used with existing VCF tools such as tabix and IGV. This feature makes the file convenient both for direct interpretation and as a starting point for further analysis.

## gvcftools

Illumina has created a full set of utilities aimed at creating and analyzing Genome VCF files. For up to date information and downloads, visit the gvcftools website at sites.google.com/site/gvcftools/home.

## Examples

The following is a segment of a VCF file following the gVCF conventions for representation of nonvariant sites and, more specifically, using gvcftools block compression and filtration levels.

In the following gVCF example, nonvariant regions are shown in normal text and variants are shown in **bold**.



NOTE

The variant lines can be extracted from a gVCF file to produce a conventional variant VCF file.

```
chr20 676337 . T . 0.00 PASS END=676401;BLOCKAVG_min30p3a
   GT:GQX:DP:DPF 0/0:143:51:0
chr20 676402 . A . 0.00 PASS END=676441;BLOCKAVG_min30p3a
   GT:GQX:DP:DPF 0/0:169:57:0
chr20 676442 . T G 287.00 PASS SNVSB=-30.5;SNVHPOL=3
   GT:GQ:GQX:DP:DPF:AD 0/1:316:287:66:1:33,33
chr20 676443 . T . 0.00 PASS END=676468;BLOCKAVG_min30p3a
   GT:GQX:DP:DPF 0/0:202:68:1
chr20 676469 . G . 0.00 PASS END=676528;BLOCKAVG_min30p3a
   GT:GQX:DP:DPF 0/0:157:53:0
chr20 676529 . T . 0.00 PASS END=676566;BLOCKAVG_min30p3a
   GT:GQX:DP:DPF 0/0:120:41:0
chr20 676567 . C . 0.00 PASS END=676574;BLOCKAVG_min30p3a
   GT:GQX:DP:DPF 0/0:120:41:0
```

```
chr20 676575 . A T 555.00 PASS SNVSB=-50.0; SNVHPOL=3
  GT:GQ:GQX:DP:DPF:AD 1/1:114:114:39:0:0,39
chr20 676576 . T . 0.00 PASS END=676625; BLOCKAVG min30p3a
  GT:GQX:DP:DPF 0/0:95:36:0
chr20 676626 . T . 0.00 PASS END=676650; BLOCKAVG min30p3a
   GT:GQX:DP:DPF 0/0:117:40:0
chr20 676651 . T . 0.00 PASS END=676698; BLOCKAVG min30p3a
  GT:GQX:DP:DPF 0/0:90:31:0
chr20 676699 . T . 0.00 PASS END=676728; BLOCKAVG min30p3a
  GT:GQX:DP:DPF 0/0:69:24:0
chr20 676729 . C . 0.00 PASS END=676783; BLOCKAVG min30p3a
  GT:GQX:DP:DPF 0/0:57:20:0
chr20 676784 . C . 0.00 PASS END=676803; BLOCKAVG min30p3a
  GT:GQX:DP:DPF 0/0:51:18:0
chr20 676804 . G A 62.00 PASS SNVSB=-7.5; SNVHPOL=2
  GT:GQ:GQX:DP:DPF:AD 0/1:95:62:17:0:11,66
chr20 676805 . C . 0.00 PASS END=676818; BLOCKAVG min30p3a
   GT:GQX:DP:DPF 0/0:48:17:0
chr20 676819 . T . 0.00 PASS END=676824; BLOCKAVG min30p3a
  GT:GQX:DP:DPF 0/0:39:14:0
chr20 676825 . A . 0.00 PASS END=676836; BLOCKAVG min30p3a
  GT:GQX:DP:DPF 0/0:30:11:0
chr20 676837 . T . 0.00 LowGQX END=676857; BLOCKAVG min30p3a
  GT:GQX:DP:DPF 0/0:21:8:0
chr20 676858 . G . 0.00 PASS END=676873; BLOCKAVG min30p3a
   GT:GQX:DP:DPF 0/0:30:11:0
```

In addition to the nonvariant and variant regions in the example, there is also 1 nonvariant region from [676837,676857] that is filtered out due to insufficient confidence that the region is homozygous reference.

#### Conventions

Any VCF file following the gVCF convention combines information on variant calls (SNVs and small-indels) with genotype and read depth information for all nonvariant positions in the reference. Because this information is integrated into a single file, distinguishing variant, reference, and no-call states for any site of interest is straightforward.

The following subsections describe the general conventions followed in any gVCF file, and provide information on the specific parameters and filters used in the Isaac workflow gVCF output.



NOTE

 $\ensuremath{\mathsf{gVCF}}$  conventions are written with the assumption that only one sample per file is being represented.

#### Interpretation

gVCFs file can be interpreted as follows:

- ▶ Fast interpretation—As a discrete classification of the genome into 'variant', 'reference', and 'no-call' loci. This classification is the simplest way to use the gVCF. The Filter fields for the gVCF file have already been set to mark uncertain calls as filtered for both variant and nonvariant positions. Simple analysis can be performed to look for all loci with a filter value of "PASS" and treat them as called.
- ▶ **Research interpretation**—As a 'statistical' genome. Additional fields, such as genotype quality, are provided for both variant and reference positions to allow the threshold

between called and uncalled sites to be varied. These fields can also be used to apply more stringent criteria to a set of loci from an initial screen.

#### **External Tools**

gVCF is written to the VCF 4.1 specifications, so any tool that is compatible with the specification (such as IGV and tabix) can use the file. However, certain tools are not appropriate if they:

- Apply algorithms to VCF files that make sense for only variants calls (as opposed to variant and nonvariant regions in the full gVCF);
- Are only computationally feasible for variant calls.

For these cases, extract the variant calls from the full gVCF file.

## Special Handling for Indel Conflicts

Sites that are "filled in" inside deletions have additional treatment.

- ▶ Heterozygous Deletions—Sites inside heterozygous deletions have haploid genotype entries (ie "0" instead of "0/0", "1" instead of "1/1"). Heterozygous SNVs are marked with the SiteConflict filter and their original genotype is left unchanged. Sites inside heterozygous deletions cannot have a genotype quality score higher than the enclosing deletion genotype quality.
- ▶ **Homozygous Deletions**—Sites inside homozygous deletions have genotype set to "." (period), and site and genotype quality are also set to "." (period).
- ▶ All Deletions—Sites inside any deletion are marked with the filters of the deletion, and more filters can be added pertaining to the site itself. These modifications reflect the idea that the enclosing indel confidence bounds the site confidence.
- ▶ Indel Conflicts—In any region where overlapping deletion evidence cannot be resolved into 2 haplotypes, all indel and set records in the region are marked with the IndelConflict filter.

Table 27 Indel Conflict Filters

ID	Type	Description	
IndelConflict	site/indel	Locus is in region with conflicting indel calls.	
SiteConflict	site	Site genotype conflicts with proximal indel call. This conflict is typically a heterozygous genotype found inside a heterozygous deletion.	

## Representation of Non-Variant Segments

This section includes the following subsections:

- Block representation using END key
- Joining nonvariant sites into a single block record
- Block sample values
- Nonvariant block implementations

## Block Representation Using END Key

Continuous nonvariant segments of the genome can be represented as single records in gVCF. These records use the standard 'END" INFO key to indicate the extent of the record. Even though the record can span multiple bases, only the first base is provided in the REF field (to reduce file size). Following is a simplified example of a nonreference block record:

##INFO=<ID=END, Number=1, Type=Integer, Description="End position of
 the variant described in this record">
#CHROM POS ID REF ALT QUAL FILTER INFO FORMAT NA19238

The example record spans positions [51845,51862].

## Joining Non-Variant Sites Into a Single Block Record

Address the following issues when joining adjacent nonvariant sites into block records:

- ▶ The criteria that allow adjacent sites to be joined into a single block record.
- ▶ The method to summarize the distribution of SAMPLE or INFO values from each site in the block record.

At any gVCF compression level, a set of sites can be joined into a block if...

- Each site is nonvariant with the same genotype call. Expected nonvariant genotype calls are { "0/0", "0", "./.", "." }.
- ▶ Each site has the same coverage state, where 'coverage state' refers to whether at least 1 read maps to the site. For example, sites with 0 coverage cannot be joined into the same block with covered sites.
- Each site has the same set of FILTER tags.
- ▶ Sites have less than a threshold fraction of nonreference allele observations compared to all observed alleles (based on AD and DP field information). This threshold is used to keep sites with high ratios of nonreference alleles from being compressed into nonvariant blocks. In the Isaac Variant Caller gVCF output, the maximum nonreference fraction is 0.2

## **Block Sample Values**

Any field provided for a block of sites, such as read depth (using the DP key), shows the minimum observed value among all sites encompassed by the block.

## Nonvariant Block Implementations

Files conforming to the gVCF conventions delineated in this document can use different criteria for creation of block records, depending on the desired trade-off between compression and nonvariant site detail. The Isaac Variant Caller provides the following blocking scheme 'min30p3a' as the nonvariant block compression scheme.

Each sample value shown for the block, such as the depth (using the DP key), is restricted to have a range where the maximum value is within 30% or 3 of the minimum. Therefore, for sample value range [x,y],  $y \le x + max(3, x*0.3)$ . This range restriction applies to all sample values written in the final block record.

## Genotype Quality for Variant and Nonvariant Sites

The gVCF file uses an adapted version of genotype quality for variant and nonvariant site filtration. This value is associated with the GQX key. The GQX value is intended to represent the minimum of Phred genotype quality {assuming the site is variant, assuming the sites is nonvariant}.

You can use this value to allow a single value to be used as the primary quality filter for both variant and nonvariant sites. Filtering on this value corresponds to a conservative assumption appropriate for applications where reference genotype calls must be determined at the same stringency as variant genotypes, for example:

- ▶ An assertion that a site is homozygous reference at  $GQX \ge 30$  is made assuming the site is variant.
- An assertion that a site is a nonreference genotype at  $GQX \ge 30$  is made assuming the site is nonvariant.

#### Filter Criteria

The gVCF FILTER description is divided into 2 sections: (1) describes filtering based on genotype quality; (2) describes all other filters.



#### NOTE

These filters are default values used in the current Isaac Variant Caller implementation. However, no set of filters or cutoff values are required for a file to conform to gVCF conventions.

The genotype quality is the primary filter for all sites in the genome. In particular, traditional discovery-based site quality values that convey confidence that the site is "anything besides the homozygous reference genotype," such as SNV quality, are not used. Instead, a site or locus is filtered based on the confidence in the reported genotype for the current sample.

The genotype quality used in gVCF is a Phred-scaled probability that the given genotype is correct. It is indicated with the FORMAT field tag GQX. Any locus where the genotype quality is below the cutoff threshold is filtered with the tag LowGQX. In addition to filtering on genotype quality, some other filters are also applied.

The gVCF output from Isaac Variant Caller includes several heuristic filters applied to the site and indel records. The filters are as follows.

Table 28 VCF Site and Indel Record Filters

VCF Filter ID	Type Description		
HAPLOID_ CONFLICT	site/indel	te/indel Locus has heterozygous genotype in a haploid region.	
HighDepth	site/indel	The locus depth is greater than 3x the mean chromosom depth.	
HighDPFRatio	site	The fraction of base calls filtered out at a site is greater than 0.3.	
HighSNVSB	site	SNV strand bias value (SNVSB) exceeds 10.	
IndelConflict	indel	The locus is in region with conflicting indel calls.	
IndelSizeFilter	indel	Indel is outside reportable size range. Insertion/Deletion range reported in VCF header.	
LowGQX	vGQX site/indel Locus GQX is less than 30 or not present.		
SiteConflict	indel	The site genotype conflicts with the proximal indel call. This call is typically a heterozygous SNV call made inside a heterozygous deletion.	
VCF Filter ID	Type	Description	
HAPLOID_ CONFLICT	site/indel	Locus has heterozygous genotype in a haploid region.	
HighDepth	site/indel	The locus depth is greater than 3x the mean chromosome depth.	
HighDPFRatio	DPFRatio Site The fraction of base calls filtered out at a site is greate 0.3.		
HighSNVSB	site SNV strand bias value (SNVSB) exceeds 10.		
IndelConflict	indel	The locus is in region with conflicting indel calls.	
IndelSizeFilter	indel	Indel is outside reportable size range. Insertion/Deletion range reported in VCF header.	
LowGQX	site/indel	Locus GQX is less than 30 or not present.	
SiteConflict	indel	The site genotype conflicts with the proximal indel call. This call is typically a heterozygous SNV call made inside a heterozygous deletion.	

# Isaac Copy Number Variant Caller

Isaac Copy Number Variant (CNV) Caller is an algorithm for calling copy number variants from a diploid sample. Most of a normal DNA sample is diploid, or having 2 copies. Isaac CNV Caller identifies regions of the sample genome that are not present, or present either one time or more than 2 times in the genome. Isaac CNV Caller scans the genome for regions having an unexpected number of short read alignments. Regions with fewer than the expected number of alignments are classified as losses. Regions having more than the expected number of alignments are classified as gains.

Isaac CNV Caller is appropriately applied to low-depth cytogenetics experiments, low-depth single-cell experiments, or whole-genome sequencing experiments. Isaac CNV Caller is not appropriate for whole exome experiments, cancer studies, or any other experiment with the following conditions:

- Most of the genome is not assumed to be diploid.
- Reads are not distributed randomly across the diploid genome.

#### Workflow

Isaac CNV Caller can be conceptually divided into 4 processes:

- ▶ Binning—Counting alignments in genomic bins.
- ▶ Cleaning—Removal of systematic biases and outliers from the counts.
- ▶ Partitioning—Partitioning the counts into homogenous regions.
- ▶ Calling—Assigning a copy number to each homogenous region.

These processes are explained in subsequent sections.

## **Binning**

The binning procedure creates genomic windows, or bins, across the genome and counts the number of observed alignments that fall into each bin. The alignments are provided in the form of a BAM file.

Isaac CNV Caller binning keeps in memory a collection of BitArrays to store observed alignments, one BitArray for each chromosome. Each BitArray length is the same as its corresponding chromosome length. As the BAM file is read in, Isaac CNV Caller records the position of the left-most base in each alignment within the chromosome-appropriate BitArray. After all alignments in the BAM file have been read, the BitArrays have a "1" wherever an alignment was observed and a "0" everywhere else.

After reading in the BAM file, a masked FASTA file is read in, one chromosome at a time. This FASTA file contains the genomic sequences that were used for alignment. Each 35-mer within this FASTA file is marked as unique or nonunique with uppercase and lowercase letters. If a 35-mer is unique, then its first nucleotide is capitalized; otherwise, it is not capitalized. For example, in the sequence:

 $\verb|acgtttaATgacgatGaacgatcagctaagaatacgacaatatcagacaa|\\$ 

#### The 35-mers marked as unique are as follows:

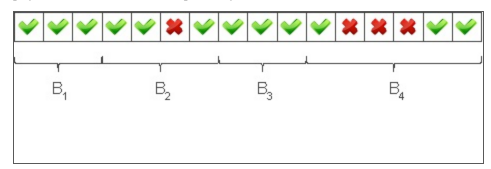
ATGACGATGAACGATCAGCTAAGAATACGACAATATGACGATGAACGATCAGCTAAGAATACGACAATATCAGACAATATCAGACAA

Isaac CNV Caller stores the genomic locations of unique 35-mers in another collection of BitArrays analogous to BitArrays used to store alignment positions. Unique positions and nonunique positions are marked with "1"s and "0"s, respectively. This marking is used as

a mask to guarantee that only alignments that start at unique 35-mer positions in the genome are used.

#### Bin Sizes

Isaac CNV Caller is initialized with 100 alignments per bin and then proceeds to compute the bin boundaries such that each bin contains the same bin size, or number of unique 35-mers. The term "bin size" refers to the number of unique genomic 35-mers per bin. Because some regions of the human genome are more repetitive than others, physical bin sizes (in genomic coordinates) are not identical. In the following example, each box is a position along the genome. Each checkmark represents a unique 35-mer while each X represents a nonunique 35-mer. The bin size in this example is 3 (3 checkmarks per bin). The physical size of each bin is not constant. B1 and B3 have a physical size of 3 but B2 and B4 have physical sizes of 4 and 6, respectively.



## Computing Bin Size

To compute bin size, the ratio of observed alignments to unique 35-mers is calculated for each autosome. The desired number of alignments per bin is then divided by the median of these ratios to yield bin size. For whole-genome sequencing, bin sizes are typically in the range of 800–1000 unique 35-mers. Correspondingly, most physical window sizes are in the 1–1.2 kb range. The advantage of this approach relative to using fixed genomic intervals is that the same number of reads map to each bin, regardless of "uniqueness" or ability to be mapped.

After bin size is computed, bins are defined as consecutive genomic windows such that each bin contains the same bin size, or number of unique 35-mers. The number of observed alignments present within the boundary of each bin is then counted from the alignment BitArrays. The GC content of each bin is also calculated. The chromosome, genomic start, genomic stop, observed counts and GC content in each bin are output to disk.

## Cleaning

The Isaac CNV Caller cleaning comprises the following 3 procedures that remove outliers and systematic biases from the count data computed in Isaac CNV Caller.

- 1 Single point outlier removal.
- 2 Physical size outlier removal.
- 3 GC content correction.

These procedures are performed on the bins produced during the Isaac CNV Caller binning process.

## Single Point Outlier Removal

This step removes individual bins that represent extreme outliers. These bins have counts that are very different from the counts present in upstream and downstream bins. Two values, a and b, are defined as to be very different when their difference is greater than expected by chance, assuming a and b come from the same underlying distribution. These values use the Chi-squared distribution, as follows:

$$\mu = 0.5a + 0.5b$$
 $\chi 2 = ((a - \mu)^2 + (b - \mu)^2) \mu - 1$ 

A value of  $\chi 2$  greater than 6.635, which is the 99th percentile of the Chi-squared distribution with 1 degree of freedom, is considered very different. If a bin count is very different from the count of both upstream and downstream neighbors, then the bin is deemed an outlier and removed.

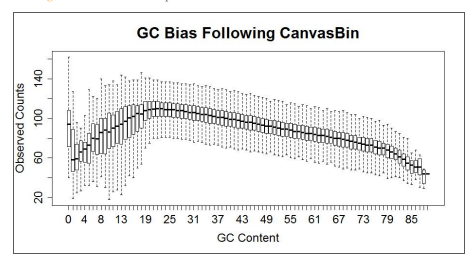
## Physical Size Outlier Removal

Bins likely do not have the same physical (genomic) size. The average for whole-genome sequencing runs might be approximately 1 kb. If the bins cover repetitive regions of the genome, some bins sizes might be several megabases in size. Example regions might include centromeres and telomeres. The counts in these regions tend to be unreliable so bins with extreme physical size are removed. Specifically, the 98th percentile of observed physical sizes is calculated and bins with sizes larger than this threshold are removed.

#### **GC Content Correction**

The main variability in bins counts is GC content. An example of the bias is represented in the following figure.

Figure 2 GC Bias Example



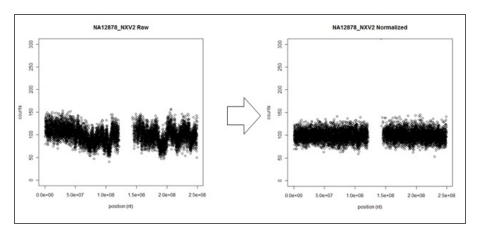
The following correction is performed:

- 1 Bins are first aggregated according to GC content, which is rounded to the nearest integer.
- 2 Second, each bin count is divided by the median count of bins having the same GC content.

Finally, this value is multiplied by the desired average count per bin (100 by default) and rounded to the nearest integer. The effect is to flatten the midpoints of the bars in the example box-and-whisker plot.

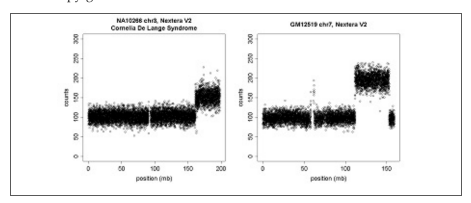
Some values for GC content have few bins so the estimate of its median is not robust. Therefore, bins are discarded when the number of bins having the same GC content is fewer than 100.

For some sample preparation schemes, GC content correction has a dramatic effect. The following figure illustrates the effect of GC content correction for a low depth sequencing experiment using the Nextera library preparation method. The figure on the left shows bins counts as a function of chromosome position before normalization. The figure on the right shows the result after GC content correction.



For whole-genome sequencing experiments, the typically median absolute deviations (MADs) are 10.3, which is close to the expected value of 10. The expected value is predicted using the Poisson model for an average count of 100 and indicates that little bias remains following GC content correction.

It is important to note that the normalization signal does not dampen signal from CNVs as shown in the following 2 figures. The figure on the left shows a chromosome known to harbor a single copy gain. The figure on the right shows chromosome known to harbor a double copy gain.



## **Partitioning**

The Isaac CNV Caller partitioning implements an algorithm for identifying regions of the genome such that their average counts are statistically different than average counts of neighboring regions. The implementation is a port of the circular binary segmentation (CBS) algorithm.

The algorithm briefly considers each chromosome as a segment. The algorithm assesses each segment and identifies the pair of bins for which the counts in the bins between them are maximally different than the counts of the rest of the bins. The statistical significance of the maximal difference is assessed via permutation testing. If the difference is statistically significant, then the procedure is applied recursively to the 2 or 3 segments created by partitioning the current segment by the identified pair of points. Input to the algorithm is the output generated by the Isaac CNV Caller cleaning algorithm.

Because of the computational complexity of the algorithm  $O(N^2)$ , the problem is divided into subchromosome problems followed by merging, in practice. Heuristics are used to speed up the permutation testing.

## Calling

The final module of the Isaac CNV Caller algorithm is to assign discrete copy numbers to each of the regions identified by the Isaac CNV Caller partitioner.

A Gaussian model is used as the default calling method. In this case, both the mean and standard deviation are estimated from the data for the diploid model and adjusted for the other copy number models. For example, if the mean,  $\mu$ , and standard deviation,  $\sigma$ , are estimated to be 100 and 15 in the diploid model, then corresponding estimates in the haploid model would be  $\mu/2$  and  $\sigma/2$ . The mean and standard deviation are estimated using the autosomal median and MAD of counts. This model is the default as it is more appropriate in cases where the spread of counts is higher than expected from the Poisson model due to unaccounted sources of variability. An example of this case is single cell sequencing experiments where whole-genome amplification is required.

Following assignment of copy number states, neighboring regions that received the same copy number call are merged into a single region.

Phred-scaled Q-scores are assigned to each region using a simple logistic function derived using array CGH data as the gold standard. The probability of a miscall is modeled as

$$p=1-(1/((1+e^{(0.5532-0.147N))})$$

Where N is the number of bins found within the nondiploid region. This probability is converted to a Q-score by

This estimate is likely conservative as it is derived from array CGH. Importantly, Q-scores are a function of number of bins, not genomic size, so they are applicable to experiments of any sequencing depth, including low-depth cytogenetics screening.

The coordinates of nondiploid regions and their Q-scores are output to a VCF file. Two filters are applied to PASS variants. First, a variant must have a Q-score of Q10 or greater. Second, a variant must be of size 10 kb, or greater.

## Isaac Structural Variant Caller

Isaac Structural Variant (SV) Caller is a structural variant caller for short sequencing reads. It can discover structural variants of any size and score these variants using both a diploid genotype model and a somatic model (when separate tumor and normal samples are specified). Structural variant discovery and scoring incorporate both paired read fragment spanning and split read evidence.

#### **Method Overview**

Isaac SV Caller works by dividing the structural variant discovery process into 2 primary steps–scanning the genome to find SV associated regions and analysis, scoring, and output of SVs found in such regions.

#### 1 Build SV association graph

In this step, the entire genome is scanned to discover evidence of possible SVs and large indels. This evidence is enumerated into a graph with edges connecting all regions of the genome that have a possible SV association. Edges can connect 2 different regions of the genome to represent evidence of a long-range association, or an edge can connect a region to itself to capture a local indel/small SV association. These associations are more general than a specific SV hypothesis, in that many SV candidates can be found on 1 edge, although typically only 1 or 2 candidates are found per edge.

#### 2 Analyze graph edges to find SVs

The second step is to analyze individual graph edges or groups of highly connected edges to discover and score SVs associated with the edges. These substeps of this process include:

- Inference of SV candidates associated with the edge.
- Attempted assembly of the SVs break-ends.
- Scoring and filtration of the SV under various biological models (currently diploid germline and somatic).
- Output to VCF.

## Capabilities

Isaac SV Caller can detect all structural variant types that are identifiable in the absence of copy number analysis and large scale de novo assembly. Detectable types are enumerated in this section.

For each structural variant and indel, Isaac SV Caller attempts to align the break-ends to base pair resolution and report the left-shifted break-end coordinate (per the VCF 4.1 SV reporting guidelines). Isaac SV Caller also reports any break-end microhomology sequence and inserted sequence between the break-ends. Often the assembly fails to provide a confident explanation of the data. In such cases, the variant is reported as IMPRECISE, and scored according to the paired-end read evidence alone.

The sequencing reads provided as input to Isaac SV Caller are expected to be from a paired-end sequencing assay that results in an inwards orientation between the 2 reads of each DNA fragment. Each read presents a read from the outer edge of the fragment insert inward.

#### **Detected Variant Classes**

Isaac SV Caller is able to detect all variation classes that can be explained as novel DNA adjacencies in the genome. Simple insertion/deletion events can be detected down to a configurable minimum size cutoff (defaulting to 51). All DNA adjacencies are classified into the following categories based on the break-end pattern:

- Deletions
- Insertions
- Inversions
- ▶ Tandem Duplications
- Interchromosomal Translocations

#### **Known Limitations**

Isaac SV Caller cannot detect the following variant types:

- Nontandem repeats/amplifications
- ▶ Large insertions—The maximum detectable size corresponds to approximately the read-pair fragment size, but note that detection power falls off to impractical levels well before this size.
  - FastTrack WGS service reports called variants that are 50–10 kb in size.
- Small inversions—The limiting size is not tested, but in theory detection falls off below ~200 bases. So-called microinversions might be detected indirectly as combined insertion/deletion variants.

More general repeat-based limitations exist for all variant types:

- ▶ Power to assemble variants to break-end resolution falls to 0 as break-end repeat length approaches the read size.
- ▶ Power to detect any break-end falls to (nearly) 0 as the break-end repeat length approaches the fragment size.
- The method cannot detect nontandem repeats.

While Isaac SV Caller classifies novel DNA-adjacencies, it does not infer the higher level constructs implied by the classification. For instance, a variant marked as a deletion by Isaac SV Caller indicates an intrachromosomal translocation with a deletion-like break-end pattern. However, there is no test of depth, b-allele frequency, or intersecting adjacencies to infer the SV type directly.

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## **BAM File FAQ**

A large volume of data represents the sequence and corresponding alignments, which are provided in BAM format. There are a few methods to convert BAM into different formats, such as FASTQ files.

#### Picard Tools FASTQ Extraction

Many pipelines start from FASTQ files.

Wall Clock Time: 3 hrs 54 min

To convert BAM files to FASTQ files using Picard tools, refer to the following example.

```
# Convert bam into read1.fastq and read2.fastq
$java -jar /picard-tools-1.110/SamToFastq.jar INPUT=Example.bam
    FASTQ=Example_R1.fastq SECOND_END_FASTQ=Example_R2.fastq
    VALIDATION_STRINGENCY=SILENT

BAM Size: 79 G
```

```
Optional arguments:
```

thread.

- ▶ RE\_REVERSE=true—Reverts the sequence to the native orientation. Otherwise, all aligned sequence is forward orientation.
- MAX\_RECORDS\_IN\_RAM=5000000—Decides the number of reads held memory and controls total memory usage.

Picard requires large amounts of memory. Picard reads data sequentially line by line from the BAM file and stores the reads in memory until both pairs of each read have been read. Memory is reset only when the reads are printed. Every read that does not have adjacent or near adjacent pairs requires more memory. Therefore, sort large BAM files when memory is a limiting factor.

For additional information about Picard Tools, see picard.sourceforge.net/command-lineoverview.shtml.

Download Picard Tools at sourceforge.net/projects/picard/files/picard-tools.

#### SAMtools Sort

SAMtools sort ensures that paired reads are next to each other, so you can save a significant amount of memory by using SAMtools to sort the BAM files by name before running Picard.

For additional information about SAMtools, see samtools.sourceforge.net/

## Reads Extraction Using SAMtools Flags

The Bam/Sam format contains a "bitwise flag" column that contains a hexadecimal, which defines the nature of that read. SAMtools allows you to easily filter on reads based on this flag. There are 12 types of such flags and using the including (-f) or the excluding (-F) option with flags from SAMtools, you can filter/extract any kind of read from the Bam/Sam file. The hexadecimal outputs are a bit hard to decipher. To convert the SAMtools flags into a human readable format, you can input the flag into picard.sourceforge.net/explain-flags.html or run the following command to output the flags in the coded string format described in the samtools manual.

```
$samtools view -X Example.bam
```

A few commonly used examples of filtering on flags are detailed below:

```
Extract all reads that are unmapped
```

```
# -f 4 = include reads which are unmapped
# command will output all the reads which are not mapped.
$samtools view -h -f 4 Example.bam
```

#### Extract reads with unmapped mates

```
# -f 8 = include reads whose mates are not mapped
# command will output all reads whose mates are not mapped.
$samtools view -h -f 8 Example.bam
```

#### Extract an unmapped read with a mapped mate

```
# -f 4 = include reads which are unmapped
# -F 8 = exclude reads whose mate is not mapped
# command outputs reads that are unmapped with the corresponding
   mate mapped
$samtools view -h -f 4 -F8 Example.bam
```

#### Extract a mapped read with an unmapped mate

```
# -f 8 = include reads whose mate is unmapped
# -F 8 = exclude all reads not mapped
# command outputs reads which are mapped with the mate is unmapped
$samtools view -h -f 8 -F4 Example.bam
```

#### Extract both reads of a pair, which are unmapped

```
#-f 12 = a combination of flag 4 and flag 8 (4+8) -> include only
   if a read is unmapped and the mate is unmapped.
# command outputs read pairs with both pairs unmapped
$samtools view -h -f 12 Example.bam
```

# Illumina FastTrack Services Annotation Pipeline

The Illumina FastTrack Services Annotation Pipeline provides variant annotation for Single Nucleotide variants (SNVs), insertions, and deletions (indels). All annotations are provided in the INFO field of [Sample\_Barcode].vcf.gz file and documented in the header.

Larger variants (CNAs, SVs) are not annotated with the full pipeline. The annotation database is queried for each of the small variants input to the pipeline. Both positional and allelic annotations can be returned for a given variant. After querying the annotation database, novel variants (variants for which no annotation exists) are then processed with VEP. If VEP does not return an annotation for the variant, it will remain unannotated.

#### **Annotation Database Sources**

The following table includes sources for the annotation databases.

Table 29 List of Annotation Database Sources

Source	Version	Release Date	
Variant Effect Predictor	72	06/01/2013	
1000 Genomes Allele Frequencies	v3, Release 20110521	04/30/2012	
ClinVar	20130905	09/05/2013	
COSMIC	65	05/28/2013	
dbSNP	137	06/16/2012	
HGNC/RefSeq Mapping	Updated daily	07/01/2013	
NHLBI Exome Variant Server	v.0.0.20 ESP6500SI-V2	06/07/2013	
phastCons	N/A	12/06/2009	

## Technical Assistance

For technical assistance, contact Illumina Technical Support.

Table 30 Illumina General Contact Information

Website	www.illumina.com
Email	techsupport@illumina.com

Table 31 Illumina Customer Support Telephone Numbers

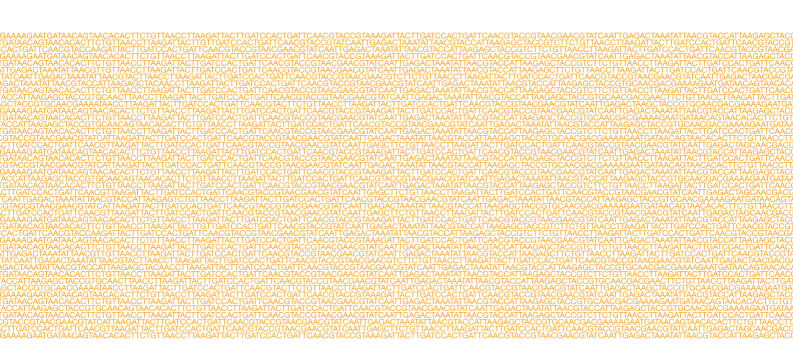
Region	Contact Number	Region	Contact Number
North America	1.800.809.4566	Italy	800.874909
Australia	1.800.775.688	Netherlands	0800.0223859
Austria	0800.296575	New Zealand	0800.451.650
Belgium	0800.81102	Norway	800.16836
Denmark	80882346	Spain	900.812168
Finland	0800.918363	Sweden	020790181
France	0800.911850	Switzerland	0800.563118
Germany	0800.180.8994	United Kingdom	0800.917.0041
Ireland	1.800.812949	Other countries	+44.1799.534000
	1.800.812949		+44.1799.53400

## Safety Data Sheets

Safety data sheets (SDSs) are available on the Illumina website at support.illumina.com/sds.html.

#### **Product Documentation**

Product documentation in PDF is available for download from the Illumina website. Go to support.illumina.com, select a product, then click **Documentation & Literature**.



Illumina 5200 Illumina Way San Diego, California 92122 U.S.A. +1.800.809.ILMN (4566) +1.858.202.4566 (outside North America) techsupport@illumina.com www.illumina.com